CLAIMS

- 1. A method for the accurate determination of van der Waals parameters for high-precision determination of crystal structures and/or energies, comprising the steps of:
 - numerically simulating at least one crystal structure based on density functional theory (DFT) calculations combined with a potential energy term representing Van der Waals interactions;
 - providing reference data containing accurate information about said at least one crystal structure;
 - defining a deviation function (F) quantifying a deviation between said reference data and said at least one simulated crystal structure;
 - fitting at least one parameter of said van der Waals potential term in such a way as to minimize said deviation function (F); and
 - obtaining the accurate van der Waals parameters from the best fit.
- A method according to claim 1, characterized in that said van der Waals potential term is defined as:

$$E_{disp} = \sum_{A,B} -f_{A,B}(r_{A,B}) \frac{C_{6,A,B}}{r_{A,B}^6}$$

wherein $f_{A,B}(r_{A,B})$ is a damping function and the sum runs over all pairs of interacting atoms, and that said fitting step comprises fitting said damping function.

3. A method according to claim 2, characterized in that said damping function is defined as

$$f_{A,B}(r) = \left(1 - \exp\left[-c\left(\frac{r}{r_{A,B}}\right)^{\frac{3}{n}}\right]\right)^{2n}$$

and that said fitting step comprises fitting the parameter $r_{A,B}$ and/or the parameter n and/or the parameter c.

 A method according to claim 2 or 3, characterized in that said fitting step furthermore comprises fitting said coefficient C_{6,A,B}.

5. A method according to any of the preceding claims, characterized in that said reference data are theoretical data obtained by Hartree-Fock calculations or Quantum Monte Carlo simulations.

- 6. A method according to any of the preceding claims, characterized in that said reference data are experimental low-temperature crystal structure data.
- 7. A method according to claim 6, characterized in that said crystal structure data are obtained by X-Ray or neutron scattering.
- 8. A method for the accurate determination of crystal structures and/or energies, comprising the steps of:
 - providing a rough estimate model of at least one crystal structure;
 - numerically simulating said at least one crystal structure based on density functional theory (DFT) calculations combined with a potential energy term representing Van der Waals interactions; and
 - obtaining said at least one crystal structure and/or its energy as a result of said numerical simulation,

characterized in that said van der Waals potential term is obtained by the method according to any of claims 1 to 7.

- 9. A method according to claim 8, characterized in that a plurality of polymorphic crystal structures are determined and ranked according to their respective energies.
- 10. A method for the efficient numerical optimization of a molecular crystal structure using an advantageous crystal coordinate system, comprising the steps of:
 - providing a starting crystal lattice described by an initial coordinate system comprising lattice parameters and atomic positions in said crystal;
 - defining a so-called natural coordinate system and representing said starting crystal lattice in said natural coordinate system, said natural coordinate system comprising:
 - first coordinates describing symmetry-allowed lattice changes and defined in such a way that changes of said first coordinates do not cause changes of the molecular geometry or a rotation of molecules with respect to each other and leave fractional coordinates of molecular centres constant;

• second coordinates describing symmetry-allowed translations of said molecules in said crystal;

- third coordinates describing symmetry-allowed rotations of said molecules in said crystal;
- fourth coordinates describing symmetry-allowed changes of the molecular geometry;
- transforming coordinates from said natural coordinate system to said initial coordinate system;
- calculating the lattice energy and energy derivatives with respect to said initial coordinate system; and
- transforming said energy derivatives from said initial coordinate system to said natural coordinate system,

wherein a minimization algorithm is used for minimizing said lattice energy with respect to said natural coordinate system.

- 11. A method for the energy ranking of polymorphic crystal structures, comprising the steps of:
 - providing rough estimate models of each of said crystal structures;
 - numerically simulating each of said crystal structures based on density functional theory (DFT) calculations combined with a potential energy term representing Van der Waals interactions.
 - obtaining accurate crystal structures and energies as a result of said numerical simulation; and
 - ranking said accurate crystal structures according to their respective acccurate energies.
- 12. The method according to claim 11, characterized in that the crystals are crystals of pharmaceutical compounds.
- 13. The method according to claim 12, characterized in that it is applied to identify the most stable polymorphic form of a pharmaceutical compound.
- 14. The method according to any of claims 11 to 13, characterized in that it uses a method for the efficient optimization of the molecular crystal structure according to claim 10.

15. A computer program product comprising computer readable code for enabling a computer to perform a method according to any of the preceding claims when said code is executed.